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Ø [21] Specific Synthesis of DNA in Vitro via Polymerase-Catalyzed Chain Reaction

By KARY B. MULLIS and FRED A. FALOONA

The sequence to be synthesized can be present initially as a discrete of the reaction will be a discrete dsDNA molecule with termini correamplified sequence or to append new sequence information to it. It is necessary that the ends of the sequence be known in sufficient detail that oligonucleotides can be synthesized which will hybridize to them, and that a small amount of the sequence be available to initiate the reaction. It is not necessary that the sequence to be synthesized enzymatically be present initially in a pure form; it can be a minor fraction of a complex molecule or it can be part of a larger molecule. In either case, the product We have devised a method whereby a nucleic acid sequence can be exponentially amplified in vitro. The same method can be used to alter the mixture, such as a segment of a single-copy gene in whole human DNA. sponding to the 5' ends of the oligomers employed.

Fig. 1. A source of DNA including the desired sequence is denatured in deoxyribonucleoside triphosphates. The oligonucleotides are compleuct of the one, when denatured, can serve as a template for the other, and The reaction products are denatured and the process is repeated until the desired amount of the 110-bp sequence bounded by the two oligonu-Synthesis of a 110-bp fragment from a larger molecule via this procedure, which we have termed polymerase chain reaction, is depicted in the presence of a large molar excess of two oligonucleotides and the four mentary to different strands of the desired sequence and at relative positions along the sequence such that the DNA polymerase extension prodvice versa. DNA polymerase is added and a reaction allowed to occur.

During the first and each subsequent reaction cycle extension of each molecule of indefinite length. These "long products" will accumulate in a linear fashion, i.e., the amount present after any number of cycles will be linearly proportional to the number of cycles. The long products thus produced will act as templates for one or the other of the oligonucleotides during subsequent cycles and extension of these oligonucleotides by polymerase will produce molecules of a specific length, in this case, 110 bases iong. These will also function as templates for one or the other of the oligonucleotides producing more 110-base molecules. Thus a chain reacoligonucleotide on the original template will produce one new ssDNA cleotides is obtained.



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oligonucleotides were PC03 and PC04 at 10 $\mu M_{\rm s}$ and dNTPs were labeled with $\alpha^{-13} P$ at 500 Ci/mol. After each synthesis cycle 10-µl aliquots were removed and these (lanes 1-10) were analyzed on a 14% polyacrylamide gel in 90 mM Tris-borate and 2.5 mM EDTA at pH 8.3 addition of $0.5~\mu g/ml$ ethidium bromide, washed with the original huffer, and photographed in UV light using a red litter. The numbers on the left margin indicate the sizes of DNA in base pairs. (B) The 110-bp fragment produced was excised from the get under UV light and of the form pinol/10 $\mu l = 0.01[(1+y)^N + yN + 1]$, where N represents the number of cycles by the manufacturer); reacted at 37° for 15 hr. PAGE was performed as above. (1) 1 µg and 24 V/cm for 2.5 hr. The completed get was soaked 20 min in the same buffer with the the incorporated 19P counted by Cerenkov radiation. An attempt to fit the data to an equation and y the fractional yield per cycle, was optimal with y = 0.619. (C) The 8- μ t aliquots from the tenth cycle of a reaction similar to the above were subjected to restriction analysis by Fig. 2. (A) Reactions were performed as in Method I. DNA target was pBR328;; BA, addition of 1 µ1 BSA (25 mg/ml) and 1 µ1 of the appropriate enzyme (undiluted, as supplied $\phi X 174/Hac IIII digest, (2) no enzyme, (3) 8 units <math>Hin \Omega$ (4) 0.5 units $Min \Omega$ (5) 2 units $Ms \Omega \Omega$ (6) 3.5 units Newl. The numbers on the left margin indicate the sizes (in base pairs) of DNA.

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tion can be sustained which will result in the accumulation of a specific 110-bp dsDNA at an exponential rate relative to the number of cycles.

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data have been fit to a simple exponential curve (Fig. 2B), which assumes remains constant over the 10 cycles. This is probably not true; however, Figure 2 demonstrates the exponential growth of the 110-bp fragment beginning with 0.1 pmol of a plasmid template. After 10 cycles of polythe precision of the available data and our present level of sophistication merase chain reaction, the target sequence was amplified 100 times. The that the fraction of template molecules successfully copied in each cycle in fully understanding the several factors involved do not seem to justify a more elaborate mathematical model. This analysis results in a calculated yield per cycle of about 62%. Amplification of this same 110-bp fragment

Fig. 1. The polymerase chain reaction amplification of a 110 bp fragment from the first exon of the human \(\beta\)-globin gene.

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Materials and Methods

dite chemistry. Synthesis and purification were performed according to Oligonucleotides were synthesized using an automated DNA synthesis machine (Biosearch, Inc., San Rafael, California) using phosphoramithe directions provided by the manufacturer.

| | | Designed to | From |
|---|--------------------------------------|--------------------|------------|
| | Oligodeoxyribonucleotides | produce | template |
| _ | FF02 CGCATTAAAGCTTATCGATG | 75 hp with 1-1-03 | pBR322 |
| | FF03 TAGGCGTATCACGAGGCCCF | | |
| | FF05 CTTCCCCATCOCTGATCTC | Son by with FFO3 | PBR 12.2 |
| _ | FF05 CCAGCAAGACGTAGCCCAGC | 1000 bp with 1-103 | pBR322 |
| | KM29 GGTTGGCCAATCTACTCCAGG | | |
| | KM30 TAACCTTGATACCAACCTGCCC | 240 bp with KM29 | Globin DNA |
| | KM38 TGGTCTCCTTAAACCTGTCTT | 268 bp with KM29 | Globin DNA |
| | KM47 AATTAATACGACTCACTATAGGGAGA. | As FF03 plus 26 hp | pBR322 |
| _ | TAGGCGTATCACGAGGCCCT | | |
| | PC03 ACACAACTGTGTTCACTAGG | | |
| | PC04 CAACITCATCCACGITCACC | 110 bp with PC03 | Globin DNA |
| | PC05 TITIGCTTCTGACACAACTGTGTTCACTAGG | | |
| _ | PC06 GCCTCACCACCACTTCATCCACGTTCACC | 130 bp with PC05 | Globin DNA |
| | PC07 CAGACACCATGGTGCACCTGACTCCTG | | |
| | PCD8 CCCCACAGGGCAGTAACGGCAGACTTCTCC | 58 bp with PC07 | Globin DNA |
| | | | |

Plasmid pBR328::BA, containing a 1.9-kb insert from the first exon of the human β -globin A allele, and pBR328:: β S, representing the β -globin Sallele, were kindly provided by R. Saiki.

Beverly, Massachusetts. Klenow fragment of Excherichia coli DNA poly-Restriction enzymes were purchased from New England Biolabs, merase was purchased from United States Biochemical Corp., Cleveland, 1 R. Saiki, S. Scharf, F. Faltoona, K. Mullis, G. Horn, H. Erlich, and N. Arnheim, Science 230, 1350 (1985).

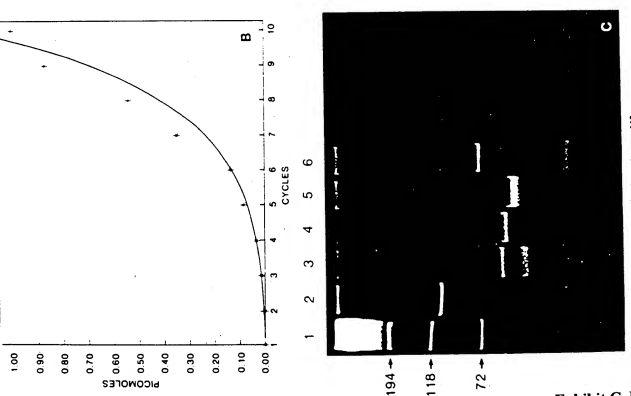


Fig. 2 (continued). See legend on p. 337.

POLYMERASE CHAIN REACTION

Ohio, and was the product of a Klenow fragment clone rather than an enzymatic cleavage of DNA polymerase 1.

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Acrylamide was from Bio-Rad Laboratories, Richmond, California: deoxyribonucleoside triphosphates were from Sigma Chemical Co., St Louis, Missouri.

NuSieve agarose was purchased from FMC Corporation. Gels were bromide. Poured into horizontal trays, the gels were ~0.5 cm thick, 10 cm long, and were run for 60-90 min at 10 V/cm submerged in the buffer described above. From 4 to 6% NuSieve agarose gels provide separations comparable to 10-15% polyacrylamide; they are considerably easier to prepared by boiling the appropriate amount of agarose in 90 mM Trisborate at pH 8.3, 2.5 mM in EDTA, and containing 0.5 µg/ml ethidium light. Prior to photography, gels are soaked in water for 20 min to remove cast and load and can be monitored while running with a hand-held UV unbound ethidium bromide.

The following method is representative of a number of PCR protocols which have been successfully utilized. Specific variations on this procedure are noted in the figure legends and several are summarized below.

Polymerase Chain Reaction: Method 1

cleotides FF02 and FF03 (3 μM) (see Diagram 1), and 150 nmol of each 7.9), 60 mM sodium acetate, 10 mM dithiothreitol, and 10 mM magnesium acetate. The solution is brought to 100° for 1 min, and is cooled to 25° for 30 sec in a waterbath. Add 1.0 μ l containing 5 units of Klenow fragment of Dissolve 0.1 pmol pBR322 (1 nM) and 300 pmol each of oligonudeoxynucleoside triphosphate (1.5 mM) in 100 µl 30 mM Tris-acetate (pH

EXTENDS <----GCTGTAGTGGCTAGCCCTTC 5' FFUS

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DIAGRAN 1. PCR model systems in pBR322

E. coli DNA polymerase I and allow the reaction to proceed for 2 min at 25°, after which the cycle of heating, cooling, adding enzyme, and reacting is repeated nine times.

Method I (Summary of Above)

Target DNA: 0.1 pmol

Oligonucleotides: 3 µM, 20-mers

Buffer: 100 µl 30 mM Tris-acetate (pH 7.9) 60 mM sodium acetate,

10 mM Magnesium acetate, and 10 mM DTT

dNTPs: 1.5 mM

Enzyme: 5 units Klenow fragment

Cycles: Number: 10

Denaturation: 100°, 1 min

Primer hybridization: 25°, 30 sec

Reaction: 25°, 2 min

Method II (Nested Primer Sets)

Target DNA: 10 μg human DNA (0.5 \times 10 $^{\circ}$ pmol)

Oligonucleotides: 2 μM , outer set: 20-mers; inner set: 27-mer and 30-mer

Buffer: 100 µl 30 mM Tris-acetate (pH 7.9), 60 mM sodium acetate, 10 mM magnesium acetate, and 10 mM DIT

dNTPs: 1.0 mM

Enzyme: 2 units Klenow fragment

primers, a 10-µl aliquot of this reaction was diluted into a Cycles: Following 20 cycles of amplification with the outer-set further 100-µl reaction mixture containing the inner-set primers and 10 more cycles were performed.

Denaturation: 100%, 1 min

Primer hybridization: 25°, 1 min

Reaction: 25°, 2 min

Method III

Target DNA: 1 μg to 20 ng human DNA (0.5 \times 10 $^{\rm h}$ to 1 \times 10 $^{\rm s}$

Buffer: 100 µl 10 mM Tris-chloride (pH 7.5), 50 mM sodium acetate, and 10 mM magnesium chloride

Oligonucleotides: 1 µM, 20-mers

pmol)

dNTPs: 1.5 mM

Enzyme: 1 unit Klenow fragment

Cycles: Number: 20-25

Denaturation: 95°, 5 min, first cycle

Primer hybridization: 30°, 2 min 95°, 2 min, subsequent cycles

Reaction: 30°, 2 min

Method IV?

Target DNA: 1 µg human DNA (0.5 × 10 ° pmol)

Oligonucleotides: 1 μM , 20–28-mers

Buffer: 100 µl 30 mM Tris-acetate (pH 7.9), 60 mM sodium acetate,

ŧ

10 mM Magnesium acetate

UNTPS: 1.5 mM

Enzyme: 1 unit Klenow fragment

Cycles: Number: 20

Denaturation: 95°, 2 min

Primer hybridization: 37°, 2 min

Reaction: 37°, 2 min

As Method IV except

Buffer: 10% DMSO added to Method 1V buffer

Cycles: Number: 27

Method VI

Target DNA: 5 ng human DNA containing target + 250 ng human DNA deleted for target, or 1 µg human DNA containing an unknown amount of HTLV-III viral DNA sequence

Oligonucleotides: 1 µM, 15-18-mers

Buffer: 100 µl 10 mM Tris-chloride (pH 7.5), 50 mM sodium chlo-

ride, and 10 mAI magnesium chloride

dNTPs: 1.5 mM

Enzyme: I unit Klenow fragment

Denaturation: 95°, 2 min Cycles: Number: 20-25

Primer hybridization: 25°, 2 min

Reaction: 25", 2 min

Specificity of the Amplification Reaction

This process has been employed to amplify DNA segments from 24 to 000 bp in length using template DNA ranging in purity from a highly

28. Schart, G. Horn, and H. Erlich, submitted for publication.

S. Kwok, D. Mack, K. Mullis, B. Poiesz, G. Ehrlich, D. Blair, A. Friedman-Kien, and J. J.

POLYMERASE CHAIN REACTION

tions the specificity of the overall reaction is intrinsically high, probably due to the requirement that two separate and coordinated priming events occur at each cycle. Beginning with purified plasmid DNA as initial template and pairs of primers intended to produce fragments in the range of 200 bp or less, homogeneous products have usually been observed. Using similar templates, but primers chosen to amplify larger fragments, longer reaction times are required and considerable production of DNA fragments other than that intended is observed (Fig. 3). These by-products are usually smaller than the intended product and can be accounted for by "mispriming" events wherein the 3' end of one of the primers interacts with a region of partial homology within the sequence of the primary product (see Diagram 2). The probability for synthesis of a by-product quence in the original reaction for two reasons. First, the concentration of purified synthetic single-stranded DNA to a totally unpurified single-copy gene in whole human DNA. Despite the low stringency of the hybridizarepresenting a subfragment of the primary product is higher than the probability for synthesis of a hy-product representing some different sethe primary product becomes relatively high during the reaction; and result in the production of a new molecule, which like the primary product complementary to it.) The synthesis of multiple DNA fragments is thus will contain two primer sites. (A primer "site" in this context would be either a region complementary to one of the primers or a region containing one of the primers, which would in successive cycles produce a sequence second, any single "mispriming" on a molecule of primary product will more likely if the intended fragment is large and the final desired concentration of the product is high. The ~225-bp by-product of the amplification of a 500-bp fragment from pBR322 depicted in Fig. 3B can be ac-

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DIAGRAM 2. Probable second priming site on pBR 02 for F101,

Sninsky, submitted for publication.

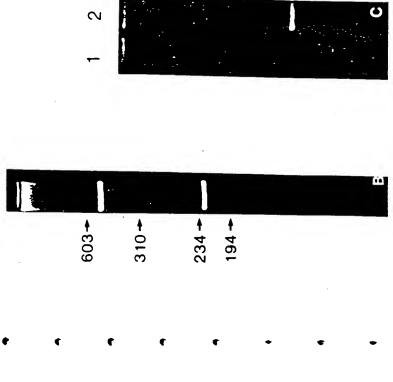


Fig. 3. (continued)

counted for by a second priming site for FF03 in which 9 out of 11 of the 3' nucleotides of FF03 find a match within the amplified product.

In Vitro Mutations

within the target sequence yields no amplified product. The numbers on the left margin

prior to the reaction. This plasmid is cut several times by Mst11 but not within the sequence to be amplified by KM29 and KM30. A similar reaction with pBR328::etaA which is cut indicate the sizes (in base pairs) of DNA. (B) Reactions were performed as in Method I. except reaction times were 20 min per cycle at 37°. Oligonucleotides were FF03 and FF05.

oligonucleotides were (1) PC'03 and PC04. (2) PC'05 and PC'06. (3) KM29 and KM38 (reaction time was 20 min), (4) KM29 and KM30; DNA target was pBR328 :: eta S digested with Msrt11

Fig. 3. (A) Reactions were performed as in Method 1. DNA target was pBR328:: β A,

Final product was rehybridized for 15 hr at 57°. Electrophoresis was on a 47° NuSieve agarose gel. The numbers on the left margin indicate the sizes (in base pairs) of DNA. (C) (1) Reactions were performed as in Method L. Oligonucleotides were FF02 and FF03. The tenth reaction cycle was terminated by freezing and an 8-µl aliquot was applied to a 4%. NuSieve agarose gel visualized with ethidium bromide. (2) Reactions were the same as in (1) except

"Mispriming" can be usefully employed to make intentional in vitro mutations or to add sequence information to one or both ends of a given sequence. A primer which is not a perfect match to the template sequence but which is nonetheless able to hybridize sufficiently to be enzymatically extended will produce a product which contains the sequence of the When this product in a subsequent cycle is template for the second primer primer rather than the corresponding sequence of the original template. the extension product produced will be a perfect match to the first primer

bp fragment, 26 nucleotides of which are not present in pBR322. The numbers on the left

that the ofigonucleotides used were FF02 and KM47, which were designed to produce a 101-

Oligonucleotides were FF03 and FF06. (2) Same as (1) except that KM47 was substituted for

FF03. The numbers on the left margin indicate the sizes (in base pairs) of DNA.

margin indicate the sizes (in base pairs) of DNA. (D) (1) Reactions were performed as in (B).

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POLYMERASE CHAIN REACTION

Fig. 3 (continued)

and an in vitro mutation will have been introduced. In further cycles this mutation will be amplified with an undiminished efficiency since no further mispaired primings are required.

can be used to insert a new sequence in the product adjacent to the template sequence being copied. In Fig. 3C, lane 2, a 26-bp T7 phage A primer which carries a noncomplementary extension on its 5' end

promoter has been appended to a 75-bp sequence from pBR322 by using sion. The procedure required less than 2 hr and produced 2 pmol of the an oligonucleotide with 20 complementary bases and a 26-base 5' extenrelatively pure 101-bp fragment from 100 fmol of pBR322. Similarly in Fig. 3D, the T7 promoter has been inserted adjacent to a 1000-bp fragment rom pBR322.

Scharf et al.,2 in order to facilitate the cloning of human genomic fragments, inserted restriction sites onto the ends of amplified sequences by the use of primers appropriately mismatched on their 5' ends.

Detection of Minute Quantities of DNA

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sequences in whole human DNA or other similarly complex mixtures of A microgram of human DNA contains 5 \times 10 $^{-19}$ moles of each singlecopy sequence. This is ~300,000 molecules. Detection of single-copy nucleic acids presents a problem which has only been successfully approached using labeled hybridization probes.

Saiki et al., by combining a PCR amplification with a labeled hybridtainty involved in determining the sequence of a single base pair change in ization probe technique, have significantly reduced the time and uncerthe human genome from only a microgram of DNA. They performed a 20. fold increase in the level of a 110-bp sequence in the first exon of the $oldsymbol{eta}_{-}$ globin gene. Once amplified the sequence was relatively simple to cycle amplification, which required less than 2 hr, and achieved a 200,000analyze.

level so as to enable visual detection via ethidium bromide staining of a We attempted to amplify the same 110-bp fragment to a slightly higher gel. For fragments in this size range. 100 fmol gives rise to a clearly visible band, thus, 0.1 aliquot of a 200,000-fold amplification of 10 μg of human DNA should be sufficient. And so it is; however, control experiments with DNA from a cell line harboring a eta-globin deletion indicated that the 110-bp fragment produced was not exclusively representative of the $oldsymbol{eta}$ globin focus. That is, fragments of ~110 bp were being amplified even though no $oldsymbol{eta}$ -globin sequences were present. On the chance that whatever was causing this "background" might not share extensive homology with eta-globin in the central 60 nucleotides of this 110-bp region, we attempted to increase the specificity of the process by introducing a second stage of amplification using a second set of primers nested within the first (see Diagram 3). By requiring four separate priming events to take place, we were thus able to amplify approximately 2,000,000-fold and readily detect a β-globin-specific product (Fig. 4).

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EXTENDS < --- TOLAMAGE GET INC 5" FIRST STACE PCR: OUTER PRIMERS AMPLIET PRIMARY FRAGMENT

5' GGLCATATAACCG--->EXTENUS

SECUND STAGE POR: INNER PRIMERS AMPLIFY SUB-FRAGMENT

EXTENDS <--- ICABGGC I ABULAL

111AGAGICCGT1****AG11CCGA11CGIG AAA1C1CAGGGCAA******TCAAGGC1AAGCAC

DIAGRAM 3. Nested primer sites, which enable a second layer of specificity. (The sequences here are only examples and have no particular significance.) The wild-type β -globin allele can be distinguished from the sickle-type product subsequent to amplification, will serve to distinguish between Ddel treatment of the DNA prior to amplification, or of the amplified allele by the presence of a site for the restriction enzyme Ddel. Thus, these two allelles.

Scharf et al.,2 beginning with I µg of human DNA and oligonucleotides 26 and 28 nucleotides in length that were designed to amplify a 240-bp region of the III.A-DQ-a gene after 27 cycles of PCR, were able to visualize the predicted fragment via ethidium staining of an agarose gel. In contrast to our results with β -globin, controls with III.A-deleted cell lines revealed that this single-stage amplification was specific for the intended

lemperature, have been forthcoming; however, the number of examples Similar amplifications of other human loci have resulted in varying ability, based on, for example, oligomer size, target size, sequence, and degrees of specificity and efficiency. No simple explanations for this variof attempted amplifications of different human sequences is still small. larget.

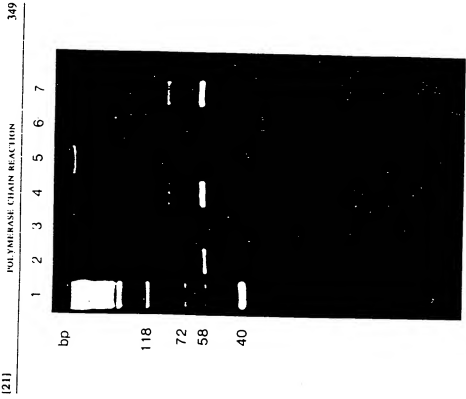


Fig. 4. Reactions were performed as in Method II, and 8 μ l aliquots (representing 80 ng with ethidium bromide. Oligonucleotides were PC03 and PC04, followed by PC07 and PC08 of unamphilied DNA) were subjected to electrophoresis on a 4% NuSieve agarose gel stained type eta-globin allele; lane (3), as in (2) but treated prior to amplification with Ddet, which cleaves the intended target and prevents amplification; lane (4), human DNA homozygous for the sickle $oldsymbol{eta}$ -globin allele treated prior to amphification with Ddel, which for this allele amplification an aliquot of the reaction in (2) was subjected to cleavage with Ddct, which (the nested set). DNA target was as follows: fane (2), human DNA homozygous for the wilddoes not cleave the intended target; lane (5), salmon sperm DNA. Following the final should convert the 58-bp wild-type product into 27- and 31-bp fragments (lane (6)); an aliquot of the reaction in (4) was similarly treated with Ddel after amplification (lane (7)). The S8-bp product from the sickle allete, as expected, contains no Pdrl site. The numbers on the left margin indicate the sizes (in base pairs) of DNA.

[2]

opmental DNA diagnostic procedures^{1,3} and in molecular cloning from genomic DNA2; it should be useful wherever increased amounts and relative purification of a particular nucleic acid sequence would be advantageous, or when alterations or additions to the ends of a sequence are The polymerase chain reaction has thus found immediate use in develrequired

cycle of thermal denaturation; in addition, it is anticipated that increasing the temperature at which the priming and polymerization reactions take merase so as to avoid the need for addition of new enzyme after each We are exploring the possibility of utilizing a heat-stable DNA polyplace will have a beneficial effect on the specificity of the amplification.

Acknowledgments

We wish to acknowledge the interest and support of Thomas White, and we would like to thank Corey Levenson, Lauri Goda, and Dragan Spasic for preparation of oligonucleotides; Randy Saiki, Stephen Scharf, Glenn Horn, Henry Erlich, Norman Arnheim, and Ed Sheldon for useful discussions regarding the amplification of human sequences; and Denise Ramirez for assistance with the manuscript.

[22] Visual Assay for Chromosome Ploidy By Douglas Koshi and and Philip Hieler

Introduction

also complex, consisting of many trans-acting factors often assembled in transmission in the yeast, Succharomyces cerevisiae, occur as infredivision requires the correct execution of a large number of biochemical reactions. The substrate in these reactions, the chromosome, is complex, exhibiting morphological and functional differentiation along its length as evidenced by the presence of specialized domains such as centromeres and telomeres. The cellular machinery that catalyzes these reactions is cuted with extremely high fidelity. For example, errors in chromosome The proper replication and segregation of chromosomes in a mitotic complicated structures, for example, the spindle. The intricate nature of the substrate and machinery apparently assure that the reactions are exequently as once per 10° cell divisions. 1.2

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process. The isolation of these mutants is not trivial because replication some transmission in the yeast, S. cerevisiae, is feasible because the fidelity of chromosome transmission exhibited by wild-type strains is much greater than the fidelity needed for viability. Thus mutations that d amatically reduce but do not destroy the fidelity of the process are viable in yeast. These mutations will include hypomorphs, leaky mutations in functions essential for chromosome transmission, or null muta-An understanding of mitotic chromosome transmission at the molecular level will only be achieved when one knows how each chromosomal ual steps of the process. This functional dissection of chromosome transmission will require genetic as well as biochemical approaches, in particular the isolation and characterization of mutants that are defective in the and segregation are essential processes and mutations which destroy domain and component of the cellular machinery functions in the individthem will be lethal to the cell. However, the genetic analysis of chromotions in functions that contribute to fidelity but are not essential for chromosome transmission.

Given that yeast mutants with altered chromosome transmission are

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¹ E. H. Hartwell, S. K. Dutcher, J. S. Wood, and B. Garvik, Revent Adv. Frust Mol. Biol. 1, 28 (1982).

t M. S. Espositio, D. T. Maleas, K. A. Bjornstad, and C. V. Brushi, Curr. Top. Genet. 6, 5

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